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ATTORNEY DOCKET NUMBER

0887-4147US2

U.S. APPLICATION NO. (If known, see 37 CFR 1.51)

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INTERNATIONAL APPLICATION NO.
PCT/US00/25136INTERNATIONAL FILING DATE
14 September 00 (14.09.00)EARLIEST PRIORITY DATE CLAIMED
14 September 99 (14.09.99)

TITLE OF INVENTION

METHOD AND SYSTEM FOR IMAGING THE DYNAMICS OF SCATTERING MEDIUM

APPLICANT(S) FOR DO/EO/US

Randall L. BARBOUR

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

1. ☒ This is a **FIRST** submission of items concerning a filing under 35 U.S.C. 371.
2. ☐ This is a **SECOND** or **SUBSEQUENT** submission of items concerning a filing under 35 U.S.C. 371.
3. ☒ This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and 21 indicated below.
4. ☒ The US has been elected by the expiration of 19 months from the priority date (Article 31).
5. ☒ A copy of the International Application as filed (35 U.S.C. 371(c)(2))
 - a. ☐ is attached hereto (required only if not communicated by the International Bureau).
 - b. ☒ has been communicated by the International Bureau.
 - c. ☐ is not required, as the application was filed in the United States Receiving Office (RO/US).
6. ☐ An English language translation of the International application as filed (35 U.S.C. 371(c)(2)).
 - a. ☐ is attached hereto.
 - b. ☐ has been previously submitted under 35 U.S.C. 154(d)(4).
7. ☒ Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3))
 - a. ☐ are attached hereto (required only if not communicated by the International Bureau).
 - b. ☐ have been communicated by the International Bureau.
 - c. ☐ have not been made; however, the time limit for making such amendments has NOT expired.
 - d. ☒ have not been made and will not be made.
8. ☐ An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
9. ☒ An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).
10. ☐ An English language translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).

Items 11. to 16. below concern document(s) or information included:

11. ☒ An Information Disclosure Statement under 37 CFR 1.97 and 1.98 with Form PTO 1498 and copy of cited reference.
12. ☐ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
13. ☐ A **FIRST** preliminary amendment.
14. ☐ A **SECOND** or **SUBSEQUENT** preliminary amendment.
15. ☐ A substitute specification.
16. ☐ A change of power of attorney and/or address letter.
17. ☐ A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 154(d)(2) with copy of Statement Under 37 CFR Section 1.821(f) and WIPO Standard ST.25 as filed with the International Bureau of WIPO.
18. ☐ A second copy of the published international application under 35 U.S.C. 154(d)(4).
19. ☐ A second copy of the English language translation of the international application under 35 U.S.C. 154(d)(4).

APPLICATION NO. (if known, see 37 C.F.R. 1.51)

INTERNATIONAL APPLICATION NO.

ATTORNEY'S DOCKET NO.

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20. ☒ Other items or information:

- Copy of published PCT Application – Publication number WO 01/20305, published on 22 March 01 with International Search Report.

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U.S. APPLICATION NO (if known, see 37 C.F.R. 1.51) 10/088190		INTERNATIONAL APPLICATION NO PCT/US00/25136		ATTORNEY'S DOCKET NUMBER 0887-4147US2	
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21. <input checked="" type="checkbox"/> The following fees are submitted: BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)): Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2) paid to USPTO and International Search Report not prepared by the EPO or JPO\$1040.00 International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO\$890.00 International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2) paid to USPTO ..\$740.00 International preliminary examination fee paid to USPTO (37 CFR 1.482) but all claims did not satisfy provisions of PCT Article 33 (1) - (4).....\$690.00 International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(1) - (4).....\$100.00 ENTER APPROPRIATE BASIC FEE AMOUNT =				CALCULATIONS PTO USE ONLY \$690.00	
Surcharge of \$130 for furnishing the oath or declaration later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(e)).					
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE		
Total claims	25 - 20 =	5	X \$18.00	\$90.00	
Independent claims	2 - 3 =	0	X \$84.00	\$ 0.00	
MULTIPLE DEPENDENT CLAIM(S) (if applicable)			+ \$280.00	\$ 0.00	
TOTAL OF ABOVE CALCULATIONS =				\$780.00	
<input checked="" type="checkbox"/> Applicant claims small entity status. See 37 C.F.R. 1.27. The fees indicated above are reduced by 1/2.				\$390.00	
SUBTOTAL =				\$390.00	
Processing fee of \$130.00 for furnishing the English translation later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(f)).				\$	
TOTAL NATIONAL FEE =				\$390.00	
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property X =				\$	
TOTAL FEES ENCLOSED =				\$390.00	
				Amount to be refunded:	\$
				charged:	\$

a. ☒ A check in the amount of \$390.00 to cover the above fees is enclosed.

b. ☐ Please charge my Deposit Account No. 13-4500, ORDER NO. _____ in the amount of \$0.00 to cover the above fees.

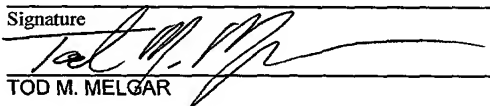
c. ☒ The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 13-4500, ORDER NO. 0887-4147US2. A duplicate copy of this sheet is enclosed.

d. ☐ Fees are to be charged to a credit card. **WARNING:** Information on this form may become public. Credit card information should not be included on this form. Provide credit card information and authorization on PTO-2038.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

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18/prts

METHOD AND SYSTEM FOR IMAGING THE DYNAMICS OF SCATTERING MEDIUM

This invention was made with U.S. Government support under contract number
5 CA-RO166184-02A, awarded by the National Cancer Institute. The U.S. Government
has certain rights in the invention.

This application claims the benefit under 35 U.S.C. §120 of prior U.S. Provisional
Patent Application Serial Nos. 60/153,926 filed September 14, 1999, entitled DYNAMIC
TOMOGRAPHY IN A SCATTERING MEDIUM and 60/154,099 filed September 15,
10 1999, entitled DYNAMIC TOMOGRAPHY IN A SCATTERING MEDIUM.

This application is related to copending application serial number "not yet
assigned", attorney docket number 0887-4147PC1, filed on the same date as this
application, entitled "SYSTEM AND METHOD FOR TOMOGRAPHIC IMAGING OF
DYNAMIC PROPERTIES OF A SCATTERING MEDIUM" by inventors R. Barbour
15 and C. Schmitz and is hereby incorporated by reference (hereinafter the "Barbour
4147PC1 application").

This application is also related to copending application serial number "not yet
assigned", attorney docket number 0887-4149PC2, filed on the same date as this
application, entitled "IMAGING OF SCATTERING MEDIA USING RELATIVE
20 DETECTOR VALUES" by inventor R. Barbour and is hereby incorporated by reference
(hereinafter the "Barbour 4149PC2 application").

Field of the Invention

The invention relates to the field of imaging in a scattering medium, and in
particular to methods of imaging the dynamics of a scattering medium.

Background

Imaging in a scattering medium relates generally to the methods and techniques for generating an image of the internal properties of a scattering medium based on the detection of scattered energy.

5 Many systems and techniques have been developed for imaging of scattering media. A typical system for imaging based on scattered energy detection includes a source for directing energy into a target medium and a plurality of detectors for measuring the scattered energy exiting the target medium at various locations with respect to the source. Based on the measured energy exiting the target medium, it is
10 possible to reconstruct an image representation of the cross-sectional scattering, absorption or other properties of the target medium. The values of the absorption and scattering properties of the medium can vary depending on the wavelength and types of energy employed as an imaging source. These values are also frequently spatially varying. These techniques permit the use of types of energy and wavelengths, such as
15 near infrared light energy, that are not suitable for projection imaging techniques, such as x-ray imaging. Thus these techniques have great potential for detecting and imaging properties of media, such as human tissue, that can not be revealed using energy sources commonly employed in projection imaging methods.

Exemplary methods and systems for imaging of scattering media are disclosed in
20 Barbour et al., U.S. Patent No. 5,137,355, entitled "Method of Imaging a Random Medium," (hereinafter the "Barbour '355 patent"), Barbour, U.S. Patent No. 6,081,322, entitled "NIR Clinical Opti-Scan System," (hereinafter the "Barbour '322 patent"), the

Barbour 4147PC1 application, the Barbour 4149PC1 application and the Barbour 4147PC2 application.

As can readily be appreciated, there are many instances where use of these techniques are highly desirable. For example, one flourishing application area is in the field of optical tomography. Optical tomography typically uses near infrared (NIR) energy as an imaging source. Contrary to imaging methods relying on the use of ionizing radiation and/or toxic/radioactive contrast agents, NIR optical tomographic imaging methods bear no risk of causing harm to the patient. The dose of optical intensity used remains far below the threshold of thermal damage and is therefore safe. In the regime of wavelength / intensity / power used, there are no effects on the tissue that accumulate with increasing light energy dose due to over-all irradiation time.

Other favorable attributes of optical tomography include the use of low-cost, potentially portable devices that employ highly integrated, economical off-the-shelf data processing electronics and semiconductor lasers (laser diodes). Such features contrast with other imaging technologies commonly used in clinical diagnosis that require large, fixed facilities such as MRI and x-ray CT imaging. Additionally, since a significant computational effort may be required for both image reconstruction and data analysis, the technology particularly gains from the exponential growth in the ratio of computing power to cost.

It is well appreciated that optical tomography has the potential to provide insights into anatomy and physiology that are unavailable from other imaging methods. For example, optical tomography, using near infrared energy, can identify the spatial variations in blood volume and blood oxygenation levels because of its sensitivity to

hemoglobin states. These measures have considerable potential value in diagnosing a broad range of disease processes that are known to influence hemoglobin states.

For example, a common feature of breast tumors, and solid tumors generally, is the occurrence of neovascularized tissue. Ultrastructurally, these tissues are highly
5 disorganized and exhibit functional abnormalities. Often the microvessels are dilated, tortuous, elongated and saccular. There is excessive branching of the vessels, including significant arterio-venous shunting as well as blind vascular endings. Aberrant vascular morphology and decreased vessel density are responsible for increase resistance to flow. The resistance to flow combined with an enlarged diffusion distance, due to the
10 expansion of the extravascular space, can lead to perfusion with hypoxemic and nutrient-deprived blood. The net effect of this state is the occurrence of substantial spatial and temporal heterogeneity in the tumor metabolic microenvironment.

Although these attributes of disease tissue are well appreciated, the availability of a suitable detection methodology able to take full advantage of these characteristics has
15 been notably lacking. An appropriate methodology would be one sensitive not only to altered hemoglobin states (i.e., localized variations in tissue blood volume and oxygenation states), but also to their dynamics under homeostatic conditions or in response to specific provocations.

A variety of methods involving imaging and non-imaging modalities are available
20 for assessing specific features of the vasculature. Detailed images of the vascular architecture involving larger vessels (> 1 mm dia.) can be provided using x-ray enhanced contrast imaging or MR angiography. These methods however are insensitive to hemoglobin states and only indirectly provide measures of altered blood flow. The latter

is well accomplished, in the case of larger vessels, using Doppler ultrasound, and for near-surface microvessels by laser Doppler measurements, but each is insensitive to variations in tissue blood volume or blood oxygenation. Ultrasound measurements are also limited by their inability to penetrate bone.

5 In principle, imaging methods based on the detection and analysis of scattered energy, such as optical tomographic methods, can provide either direct or indirect measures of all of these parameters. However, the known methods and systems have several shortcomings. First, known methods and systems provide images having low contrast and resolution. Second, these methods and systems do not image the dynamic
10 properties of highly scattering media. Third, these methods and systems require accurate calibration and are susceptible to errors. There are several reasons for these problems with known systems and methods. These reasons relate principally to how measurements are performed and how measurement data is analyzed.

For example, when imaging human tissue, the natural occurrence of vascular
15 frequencies arising from cardiac, respiratory and vasomotor activity, produces time variations in, for example, the absorption properties of tissue due to changes in tissue blood volume. Significantly, the process of vasomotion, perfusion first in one region, then another, can be expected to produce spatially convolved images should data be collected on a time frame that is long compared to the reciprocal of the frequency of these
20 processes. Thus methods that collect time-averaged data will predictably yield images whose contrast and resolution are degraded by such variability.

Also influencing the quality of reconstructed images, is the approach used to analyze the acquired data. Many of the known imaging schemes consider, in some

manner, the comparison of measured values to predicted values. Typically, these methods, including that described in the Barbour '355 patent, employ numerical methods that seek to minimize the difference between sets of measured and predicted values, and in doing so seek to provide improved estimates of the properties of the unknown target medium. These analysis schemes, referred to as model based methods, assume equivalency in the efficiency of measured detector values and computed predicted values. Although the derivation of accurate estimates of the efficiency of measured responses is possible in principle, in practice, the natural plasticity of tissue, its mainly arbitrary shape and variable composition and noted variability in hemoglobin levels, all serve to confound efforts to devise practical methods that provide reliable estimates.

In addition, the physics of light transport in highly scattering media, such as tissue, imposes further practical constraints that relate to the method adopted for data analysis. At issue is the required accuracy of assumptions made in order to generate predicted detector values, especially those adopted for the initial estimate. These assumptions are commonly referred to as the "initial guess". Small errors in the initial guess of optical properties of the reference medium can lead to large errors in the computed detector values. One consequence of this can be the severe corruption of the information content of the data vectors leading to artifact-laden images. Adding to the mentioned uncertainties is the well-known property of reconstruction methods that employ linear operators regarding their sensitivity to undetermined data sets (i.e., insufficient amount of collected data) and measurements based on restricted views (e.g., backscatter only, transmission only). The net effect of these limitations can render

solutions to image recovery problems of this type overly sensitive to the influence of experimental noise (i.e., ill-conditioned), provide nonunique solutions (ill-posed) or both.

These concerns are well appreciated by those skilled in the art of image reconstruction methods. It is also well appreciated that, in general, there are no simple or well-defined methods that can be universally applied to overcome the noted limitations. In this regard, specification of suitable conditions that can satisfactorily deal with the noted concerns is an art whose successful implementation requires considerable skill.

In addition to the need to provide stable solutions to the image reconstruction problem, consideration of the information content of the reconstructed image has considerable importance. As presently practiced, the method of optical tomography considers the evaluation of static states or employs time averaging methods to minimize the influence of signal instability originating from tissue dynamics.

The goal of these studies is to provide image maps that define spatial variations in the optical properties of tissue (usually, absorption and scatter) from which may be derived, for instance, estimates of spatial varying hemoglobin states. It is understood however, that the latter is fundamentally governed by dynamic processes whose details have the potential to reveal a wealth of information regarding functional features of the vasculature, in particular as it relates to its interaction with surrounding tissue.

The ability to measure dynamic processes of a medium can reveal information that is unobservable from static or time-averaged measures. In the case of physiological systems, the form of the dynamic process has added significance. For instance, it is well understood that many time varying processing have an underlying nonlinear character.

Nonlinear dynamic processes, in biological systems, are often chaotic and exhibit the characteristic feature of sensitivity to initial conditions.

The existence of such behavior has important implications in the understanding of disease processes and well as for the approaches taken for therapy. For instance, the approach needed to control a chaotic system is quite different from that for a linear system, wherein the system response is proportional to the magnitude of the input stimulus. Thus it has been proposed that more effective therapies can be realized from a series of well-timed perturbations rather than from the standard approach of applying a constant stimulus, the method commonly used in many pharmacological interventions. Also of interest, and related to this, is the seemingly general finding that the occurrence of chaotic behavior in physiological systems is a sign of health and its absence is a sign of disease.

For instance, it is known that heart rate variability is chaotic. Significantly, loss of this signature with the appearance of periodic oscillations is among the strongest predictors of sudden cardiac death. A similar phenomenology has been observed in infants who succumb to sudden infant death syndrome. In this case, the normally chaotic respiratory rate becomes periodic prior to the fatal incident. Similarly, during epileptic seizures, electroencephalographic recordings exhibit a transition from chaotic to periodic activity.

Presently, the capacity to monitor dynamic behavior in vascular structures is limited principally to near surface measures using laser Doppler methods. Measures of the time variability of the vascular caliber and flow motion for larger vessels is possible using Duplex ultrasound. However, these measures are insensitive to the activity of the

microvasculature, do not provide for full cross-sectional views, and are not sensitive to the dynamics of hemoglobin states.

Although optical methods, such as Laser Doppler, pulse oximetry, photoplethysmography and the like, can be used to monitor dynamic states of the vasulature, none are capable of providing such measures in the form of a cross-sectional image, especially in the case of large tissue structures. Moreover, known optical methods for cross-sectional imaging of the properties of a scattering media have not been used to derive dynamic measures of these states, and are plagued with a host of technical limitations, such as low contrast, low resolution images, prolonged computing times, excessive sensitivity to errors in initial estimates. These limitations render it unlikely that useful information regarding dynamic states could be derived from these known methods.

Overcoming the indicated drawbacks and concerns is critical for widespread practical implementation of optical tomography as a diagnostic tool because (1) improved contrast and resolution are essential to feature identification and visualization, (2) a static (snapshot) or time averaged image of a time evolving property does not provide discovery of the physiological dynamic processes and (3) measures of dynamic processes can yield critical information needed for improved diagnostic methods and therapies.

For the foregoing reasons, there is a need for a method of improving the contrast and resolution of reconstructed images. There is also a need for a method of imaging dynamic properties of dense scattering media, especially as it relates to dynamic properties of vascular states in large tissue structures as revealed by time variation in hemoglobin states. There is yet a further need for a method that can provide dynamic

images without undue reliance on complex calibration schemes or computationally intensive numerical methods.

SUMMARY OF THE INVENTION

The present invention satisfies these needs by providing a method for (1) enhancing resolution and contrast of various dynamic features of the medium being imaged, (2) imaging the dynamic properties of the target medium, and (3) generating images of the dynamic properties using techniques that reduce the need for undue system calibration and produce more stable solutions with the reconstruction algorithm.

It is one object of the invention to generate a map of the dynamic properties of a scattering medium in a cross-sectional view from a time series of collected data measurements. The measurements are obtained by directing energy into the target during a period of time and measuring the energy emerging from the target during the period of time, whereby a time series of measured energy is collected.

It is another aspect of the invention to generate a map of the cross-sectional dynamic properties of the target medium in a cross-sectional view from a time series of images of the cross-sectional properties of the target medium, wherein the dynamic properties are extracted from the time series of images using time series analysis methods.

It is a further aspect of the invention to generate a map of the cross-sectional dynamic properties of the target medium from the time series of measurements at each detector, wherein the dynamic properties are extracted from the time series of measurements at each detector using time series analysis methods.

It is a further aspect of the invention to construct a map of the dynamic properties of the medium, wherein the measured energy is processed using a modified perturbation formulation referred to as the normalized difference method, based on the radiation transport equation, using relative energy measurements. The relative energy measurements being, for example, the relative difference between an instantaneous measure and a time average mean of measures.

BRIEF DESCRIPTION OF THE FIGURES

For a better understanding of the invention, together with the various features and advantages thereof, reference should be made to the following detailed description of the preferred embodiments and to the accompanying drawings wherein:

FIG. 1 is a schematic illustration of an exemplary imaging system;

FIG. 2A is an illustration of a mechanism used to rhythmically inflate two balloons;

FIG. 2B is a plan view of the illustration of FIG. 2A;

FIG. 2C is an illustration of the geometric arrangement of optical fibers;

FIG. 3 is a reconstructed image of the cross-sectional properties of a target medium shown in FIG. 2A at an instant in time;

FIG. 4 is a cross-sectional map of the 0.2 Hz component of the amplitude of the image sequence's DFT for the target medium shown in FIG. 2;

FIG. 5 is a cross-sectional map of the 0.2 Hz component of the phase of the image sequence's DFT for the target medium shown in FIG. 2 with no phase difference between the oscillating balloons;

FIG. 6 is a cross-sectional map of the 0.2 Hz component of the phase of the image sequence's DFT, 180° phase difference between oscillating balloons;

FIG. 7A is a schematic illustration of apparatus with two-balloons;

FIG. 7B is a series of reconstructed cross-sectional images from DFT analysis of
5 image time series at 0.1 Hz and 0.24 Hz;

FIG. 8 is a cross-sectional map of ratio of FT amplitude of cardiac to respiratory frequency from forearm study;

FIG. 9 is an reconstructed MR image of forearm, with identified anatomical structures;

10 FIG. 10 is an overlay of images shown in FIGS. 8 and 9;

FIG. 11A is a graph of the CSD spectrum amplitude at position (12,19);

FIG. 11B is a graph of the CSD spectrum amplitude at position (11,10);

FIG. 11C is a graph of the CSD spectrum amplitude at position (18,27);

FIG. 12 is a cross-sectional map of amplitude, $\times 10^4$, of the DFT at the finger-flex
15 frequency;

FIG. 13 is an overlay of the map shown in FIG. 12 and an anatomical image (MRI) of the same tissue structures;

FIG. 14 is a graph of the temporal variations in μ_a (absorption coefficient) for pixels located within the involved muscle groups;

FIG. 15 is a graph of the time-dependent intensity variations for forearm measurements observed for a detector positions opposite the source;

FIG. 16 is a cross-sectional map of the correlation dimension, ν , computed, in each pixel, from time series of reconstructed images of μ_a (810 nm) obtained from
5 representative detector data shown in FIG. 15;

FIG. 17 is a cross-sectional map of the correlation dimension, ν , computed in each pixel from time series of reconstructed images of μ_a (810 nm) obtained from measurements performed on a solid white Delrin[®] rod;

FIG. 18 is a cross-sectional map of λ_1 (largest value of Lyapunov exponent) in
10 each pixel derived from the same time series of reconstructed μ_a (810 nm) images as used to generate ν map shown in FIG. 16; and

FIG. 19 is a cross-sectional comparison map of the λ_1 map in FIG. 18 to corresponding results obtained from surrogate data sets.

DETAILED DESCRIPTION OF THE INVENTION

15 Introduction

The present invention is built upon known basic approaches to tomographic imaging of a scattering medium, by enabling the investigation of the spatio temporal dynamics of a target medium. This enhancement represents a fundamentally new imaging modality with broad based applications such as in industrial processes involving
20 preparation of powdered mixtures, imaging subsurface turbulence in murky waters and

throughout clinical medicine both as a highly sensitive diagnostic imaging tool, as well as for use in treatment planning, therapeutic monitoring and follow-up.

For example, the present invention recognizes that the occurrence of dynamic behavior in human tissue lends the collected data and reconstructed images to evaluation using time series analysis methods, including pattern recognition schemes. Dynamic behavior is known to occur in vascular structures in response to cardiac, respiratory, vasomotor and local metabolic influences. Because optical tomography, based on near infrared measurements is sensitive to hemoglobin, which is nearly always restricted to the vascular compartment time variations in the measured signal, can be used to reveal the vascular frequencies. Although, similar information can be obtained by photoplethysmography, it is not available for large tissue structures nor in an cross-sectional imaging modality.

Detection of these temporal signatures/dynamic behaviors in a cross-sectional view provides a wealth of new information. For instance, the presence (or absence) of a cardiac frequency with respect to known anatomic landmarks (e.g., major arteries) provides a measure of the patency of localized arterial flow. Low-amplitude signals might suggest the presence of arterial stenosis. In addition, it is well appreciated that blood flow is under autonomic control. The absence or attenuation in amplitude of the respiratory or vasomotor frequencies might suggest an underlying peripheral neuropathy.

Further insight concerning the presence or absence of basic control mechanisms can be obtained from time-frequency analysis methods. For instance, the low-frequency vasomotor response in the major arteries of the forearm are nearly synchronous, both ipsilaterally and contralaterally, and under tight autonomic control. Disorganization of

this control might suggest the presence of pathology. In fact, loss of control mechanisms is a well known hallmark of neoplastic tissues. The nearly total absence of the vasomotor response in tumor tissue is of particular interest.

Still another factor motivating dynamic measures is an appreciation of the
5 information content available from the study of homeostatic challenges. It is well appreciated that the altered vascular architecture in tumors leads to a condition of sluggish perfusion. The existence of such states maybe be clearly revealed by induction of a homeostatic challenge. This could take on several forms. For example, simple manipulations such as deep breathing, response to cold, or mild venous congestion
10 imposed by inflation of a pressure cuff, all have been shown to produce large amplitude local variations in the optical signals attributable to tissue blood volume in the forearm. Extension of these measures to the breast and other tissue could reveal the presence of tumors with high specificity by virtue of an altered hemodynamic response.

There are other important implications to be considered concerning the value of
15 measuring the time-varying properties of hemoglobin states in a cross-sectional view. For instance, the details of the measured dynamics at the various vascular frequencies need not be the same for measures of blood volume and blood oxygenation. In addition, given the known heterogeneity in tissue perfusion, even in healthy tissue, whatever dynamic features do exist will themselves be spatially varying. Moreover, altered
20 dynamics in blood volume or blood oxygenation indicate altered local metabolic states, the existence of more central control deficits (e.g., autonomic, cardiac, respiratory), or both. Hence these measures can provide an assessment of an integrated physiological state, and also possess important features pertaining to tissue/vascular coupling. In fact,

all of the spatiotemporal features attributable to hemoglobin states that do exist will likely respond to a host of pharmacological agents and other treatment modalities. Such measures could serve to identify desired or undesired responses to therapy.

Although these examples relate to the measure of hemoglobin states, it will be appreciated by one skilled in the art that there are a plethora of dynamic features that may be observable using tomographic measures in scattering media in human tissue as well as other scattering media, such as murky water, foggy atmospheres and the like. For instance nerve activation and muscle movement can also be measured owing to changes in scattering properties of tissue. In addition, various exogenous agents that produce contrast in absorption, scatter or fluorescence can be introduced to observe dynamic states of tissue.

The present invention includes new data collection schemes and image analysis methods from which are extracted maps that reveal spatial temporal signatures of the target medium. These methods can include, linear and non-linear time series analysis methods as well as techniques for pattern recognition.

There are three principal elements to practical dynamic imaging. The first element is the use of a fast, parallel, multi-channel acquisition system. For example, a system for use in dynamic optical tomographic imaging of tissue is described briefly below and in further detail in the copending Barbour 4147PC1 application. The second element is to evaluate the acquired tomographic data to produce a cross-sectional image that reveals perturbations of the reconstructed optical coefficients from, for example, a time-averaged mean. This perturbation method is described briefly below and in further detail in the Barbour 4149PC2 application. The third element is to collect a time series of

data and subject either the time series of measured data or a time series of reconstructed images from the measure data to analysis. This can include use of various linear and nonlinear time-series analysis methods, and related techniques such as pattern recognition, to extract features identifying dynamic behavior in the properties of the target medium. These methods are discussed in further detail below.

The System

Numerous imaging systems such as those disclosed in the Barbour '355 patent, and the Barbour '322 patent have been developed for use in imaging of a scattering medium which are hereby incorporated by reference.

10 A system providing high speed data capture of one or more wavelengths simultaneous is disclosed in the Barbour 4147PC1 application. This system is capable of capturing multiple wavelength data at rates up to 150 Hz and enables the reconstruction of cross-sectional images of real-time events associated with vascular reactivity in a variety of tissue structures (e.g., limbs, breast, head and neck). Fast data collection methods are particularly useful because there are many disease states with specific influences on the spatial-dynamic properties of vascular responses in hemoglobin states.

15 A schematic illustration of an exemplary optical system for fast data collection is shown in FIG. 1. This system includes a computer 102, energy sources 104, 106, a source demultiplexer 108, an imaging head 110, detectors 112 and a data acquisition board 114.

A target 116 placed in the imaging head 110 is exposed to optical energy from sources 104, 106. The optical energy originating from energy sources 104, 106, is combined by beam splitter 118 and is delivered to source demultiplexer 108. The source

demultiplexer 108 is controlled by computer 102 to direct the optical energy to source fibers 120 sequentially. Although two energy sources 104, 106 are shown in this embodiment, an additional number of energy sources, each having different wavelengths can be employed. Moreover a single variable wavelength energy source can be

5 implemented such as Ti-Sapphire laser, dye lasers and the like.

Each source fiber 120 carries the optical energy from the demultiplexer 108 to the imaging head 110 where the optical energy is directed into the target 116. The imaging head 110 contains a plurality of source fibers 120 and detector fibers 122 for transmitting and receiving light energy, respectively. Each source fiber 120 forms a source detector

10 pair with each detector fiber 122 in the imaging head 110 to create a plurality of source detector pairs. The optical energy entering the target 116 at one location is scattered and may emerge at any location around the target 116. The emerging optical energy is collected by detector fibers 122 mounted in the imaging head 110.

The detector fibers 122 carry the emerging energy to detectors 112. The detectors

15 112 measure the energy density of the collected optical energy and generate a corresponding signal. The detectors may comprise any known photodetector such as avalanche photodiodes (APD), silicon PIN photodiodes, silicon photodiodes (SiPD), charged couple devices (CCD), charge inductive device (CID), photomultiplier tubes (PMT), multi-channel plate (MCP) and the like. The energy density is, for situations in

20 which propagating energy is any type of electromagnetic radiation, equivalent to the intensity of said radiation.

The data acquisition board 114 receives the signal, separates it by wavelength, and samples and holds the separated signals for deliver to computer 102. The computer 102 in turn reads and stores the signal for use in image reconstruction and other analysis.

Although the above described systems uses DC measurement techniques with an
5 near infrared energy source, it will be appreciated by those skilled in the art that similar systems can be implemented using other measurement techniques, such as time resolved measurements and frequency domain methods. In addition measures of acoustic signals produced in response to optical absorption (i.e., the photo acoustic effect) as well is use of acoustic sources can be similarly implemented.

10 **Data Collection, Data Analysis and Image Reconstruction**

In prior methods for imaging of scattering media, measured data was collected for a target medium and an image of the cross-sectional properties of the target was generated based on the measured data. The present invention improves on this basic process by (1) collecting a time series of measured data and (2) analyzing either the raw
15 measured data or a time series of reconstructed images to extract information relating to the dynamics of the target medium.

Data Collection

As discussed above, previously known systems and techniques sought to collect either a time averaged measure thereby minimizing the effect of dynamic behavior or to
20 collect a snapshot of the target. Both of these techniques fail to recognize the value of information contained in the time domain.

The present invention recognizes the value in measuring dynamic properties in highly scattering media and provides methods for collecting and processing the measured data to generate images of dynamic properties. These methods rely on the high speed collection of data using a system such as that described above so that a time series of data sets can be collected in the time domain. The time series analysis techniques, discussed further below, can then be used to extract the dynamic properties of the target from either the measured raw data directly or from a time series of images reconstructed from the measured data.

Data is collected in the present invention using high speed imaging systems such as the system illustrated in FIG. 1. By way of example, referring to FIG. 1, a measured data set is obtained by delivering optical energy to each of the source fibers 120 sequentially and measuring the emerging optical energy at each detector fiber 122 in parallel for each sequential source location. This process is repeated over a period of time so that a time series of data sets are generated, each data set represent a complete set of data (e.g., detector reading for each source position) for reconstruction of a cross-sectional image at an instant in time.

Reconstruction

As discussed above, the time series analysis techniques may be employed prior to image reconstruction (i.e., on the raw measured data) or after image reconstruction (i.e., on the time series of reconstructed images). Where time series analysis methods are applied to raw measured data, image reconstruction will follow based on the processed and extracted dynamic information. For example, one method may compute the discrete

Fourier transform of the measured data and then reconstruct images based on the Fourier coefficients at, for instance, a selected frequency.

Of the known reconstruction methods, one group of methods often adopted to analyze the measured data are perturbation methods. Exemplary techniques are disclosed in detail in the Barbour '355 patent and the Barbour 4149PC2 application. Briefly, in model-based techniques a perturbation formulation based on the radiation transport equation, or its approximation the diffusion equation, is generated. The radiation transport equation is a mathematical expression describing the propagation of a particle through a medium from a source location to a detector location as a function of the properties of the medium. The perturbation formulation of this equation relates a perturbation in the measure energy density at a detector to a corresponding perturbation in the properties of the medium.

The standard perturbation equation has the following forms:

$$W \cdot \delta x = \delta I \quad (1)$$

In equation (1), δI is a vector of source-detector pair intensity differences between the measured target medium and the known reference medium (i.e., $\delta I = I - I_r$). W is the weight matrix describing the influence that each volume element ("voxel") of the reference medium has on energy traveling from each source to each detector, for all source-detector pairs. The volume elements are formed by dividing a slice of the reference medium into an imaginary grid of contiguous, non-overlapping pieces. Physically, the weight matrix contains the first order partial derivatives of the detector responses with respect to the optical coefficients of each volume element of the reference medium. δx is the vector of differences between the known optical properties (e.g.,

absorption and scattering coefficients) of each volume element of the reference medium and the corresponding unknown optical properties of each volume element of the target medium.

A modification to this standard perturbation equation is presented and disclosed in the Barbour 4149PC2 application. This modified perturbation equation employs relative measurements of energy at the target detectors (e.g., the relative difference between an instantaneous measure and a time averaged mean of measures at a detector). This method improves the stability of the solution to the perturbation equation and reduces sensitivity of the equation to the selected reference medium.

The modified standard perturbation equation replaces δI in the standard perturbation equation with a proportionate relative difference between two measured values multiplied by a reference term of the required units as set forth in the equation (2) below:

$$\delta I_r = \left(\frac{I - I_0}{I_0} \right) \cdot I_r \quad (2)$$

In equation (2), I_r is the computed detector reading corresponding to a source-detector pair of a selected reference medium, and I and I_0 represent two data measurements for a corresponding source-detector pair on one or more targets (e.g., background vs. target, or time-averaged mean vs. a specific time point, etc.). The resultant term δI_r therefore represents a proportionate relative difference between two sets of measured data. This modified equation limits the effects of modeling errors and minimize ill-conditioning of the inverse problem while retaining the correct units in the solution.

The corresponding perturbation equation using this modified term is:

$$W_r \cdot \delta x = \delta I_r \quad (3)$$

In equation (3) W_r and δx are the same as W and δx in equation (1). Referring to equations (2) and (3), it can be seen that in the limit, when I_r equals to I_0 , this equation reduces to the standard perturbation formulation shown in equation (1). This formulation represents the Born approximation formulation of the modified perturbation equation. A similar expression may be written for the Rytov approximation in the following form:

$$\ell_n \left(\frac{I}{I_0} \right) = \frac{W_r}{I_r} \delta X \quad (4)$$

The inventive operation performed by equation (2) amounts to a projection procedure by which the information content of a measured proportionate relative data vector is projected onto a selected reference medium. Apart from simplifying measurement requirements, the method represented by equations (3) and (4) also reduces susceptibility of the perturbation equation to boundary and optical property variation between the target and the reference medium. Additionally, iterative solutions of equations (3) and (4) can be easily implemented. In principle, the techniques used in the modified perturbation equation can be used to evaluate any measured proportionate relative quantity to a reference quantity, for example acoustic signals.

Time Series Analysis

As discussed above, time series analysis methods may be applied to either the time series of measurement data or time series of reconstructed images. Applying these methods, the present invention enables investigation of the temporal spatial dynamics of a target medium.

Time series analysis techniques are well-known to provide measures of the frequency structure and time correlation of time varying processes. In addition, hybrid methods such as time frequency analysis (e.g., short time Fourier transform, wavelet analysis) can be used to reveal more complex behaviors in physiological systems such as

5 the known frequency modulation of the cardiac rate upon respiration. Other linear methods can be applied to time domain data such as principle component analysis to reveal the time evolution of specific processes in image maps. Practical examples of these methods as shown in FIGS. 2-19 can be used to identify, in the case of frequency analysis, specific features of the vascular tree, in particular, in the case of imaging human

10 tissue, the major arteries in the forearm. This is possible because of the known structure dependent frequency response of the vasculature (e.g., major arteries beat at a cardiac frequency). Further, time correlation methods may be employed to isolate anatomical landmarks having known phase relationships (e.g., action of antagonistic muscle groups). Additionally, similar procedures can be used on the resultant spatial map to produce the

15 relevant orthogonal information associated with these functions (i.e., maps of the amplitude or phase vs. frequency, correlation vs. time lag, etc.). In addition, complex behaviors in the form of nonlinear chaotic activity is often observable in measures of physiological systems. As is shown in FIGS. 15-19, the described invention is capable of detecting such signatures in a cross-sectional view, in particular as it relates to chaotic

20 behavior of the vasculature as measured by time variations in hemoglobin states. While such behavior has been observed in surface measurements of tissue, with the exception of the instant invention, no other imaging technique is capable of generating equivalent information.

Experimental Validation

The following discussion and some of the above examples presents results validating the methods and advantages of the present inventions. These examples are presented merely as an illustration of the benefits of the present invention and in

- 5 particular of the ability of dynamic optical tomography to provide high contrast images of time varying features in dense scattering media.

Contrast and Resolution Enhancement

- As disclosed above, application of these time-series analysis methods can extract significantly greater contrast and resolution from the optical imaging data. Image
- 10 resolution and contrast are important factors for any imaging method. Experience with optical tomography has shown that recovered images have low resolution and contrast. As an example, referring to FIG. 2, tomographic data was collected from a laboratory vessel measuring approximately 8 cm in diameter and containing four balloons 10-1 through 10-3, two of which are static and the other two were modulated at a fixed
- 15 frequency, with a relative phase of either 0° or 180° . The balloons 10 were suspended in a background scattering medium 12 consisting of 2% (v/v) Intralipid and were filled with a dilute hemoglobin solution (50 μ M). The balloons 10 were arranged in the vessel 14 as shown in FIGS. 2A and 2B. An example of a reconstructed cross-sectional image of absorption obtained at a single time point is shown in FIG. 3. Inspection of FIG. 3 shows
- 20 that whereas the four objects structure is detectable, the individual balloons are not well resolved, and the recovered contrast is poor. In contrast to this image quality, the map shown in FIG. 4, revealing the discrete Fourier transform (DFT) amplitude at the 0.2 Hz modulation frequency has nearly 100 fold increased contrast. Also evident is that the

spatial resolution of the DFT map is considerably improved compared to the maps shown in FIG. 3. The phase portions of the DFT's at 0.2 Hz for the in-phase and the 180°-out-of-phase cases are shown in FIGS. 5 and 6. The phase relations between the two modulated balloons is correctly recovered in each case.

5 A similar experiment resolving internal structures based on differences in modulation frequency is shown in FIGS. 7A and 7B. In FIG. 7A two balloons were present, one modulated at 0.1 Hz, the other at 0.24 Hz. Inspection of the reconstructed images shown in FIG. 7B, at the corresponding modulation frequencies shows complete spatio-temporal resolution.

10

Dynamic Physiological Imaging Studies on the Human Forearm

Following a protocol similar to that used to investigate dynamic behavior in the laboratory vessel, the information retrievable from dynamic studies performed on the human forearm at rest and in response to specific provocations are also explored.

15

Arm at Rest.

The natural occurrence of vascular frequencies due to respiratory and cardiac activity can be exploited to produce a spatial map revealing the presence of different functional components of the vascular tree. FIG. 8 shows a map of the logarithm of the ratio of the computed DFT amplitudes at the cardiac and respiratory frequencies obtained from a times-series of measurements on the forearm at rest. FIG. 9 is a representative MR image in the same region of the forearm. An overlay of the two maps having the same orientation is shown in FIG. 10. Inspection reveals that in the vicinity of the radial

1, interosseous artery 3 and ulnar artery 5, the ratio of the DFT amplitudes (cardiac to respiratory) is nearly ten times larger than it is in other regions. The radius 2 and ulna 4 are also shown. This response can be seen more clearly in FIG. 11A-11C, which shows the amplitude of the cross-spectral density (CSD) between a surface detector and specific locations in the image. The particular spectra shown were obtained from points in the image corresponding to locations in the flexor digitorum superficialis muscle, and points near the radial and interosseous arteries. Inspection clearly reveals that in muscle, which contains a dense matrix of microvascular structures, the dominant signal coincides with the respiratory frequency. In contrast, we observe a dominant cardiac frequency in the vicinity of the major arteries. These findings coincide well with the known hemodynamic behavior of the different structures in the vascular tree.

Finger-Flex Study.

In this study we explored further our ability to measure dynamic behavior in highly scattering media, such as the human forearm, by examining a time series of images obtained from measurements performed on a subject while conducting a finger-flex exercise. Finger flexing involves the action of so-called antagonistic muscle groups that are located on opposite sides of the forearm, specifically, the flexor digitorum superficialis on the ventral side and the extensor digitorum on the dorsal side. Results shown in FIG. 12 show a map of the amplitude of the DFT at the finger-flexing frequency (~ 0.25 Hz). FIG. 13 shows an overlay of this image onto an MR image of the same forearm in the same orientation. Inspection reveals that positions of maximum amplitude for finger-flexing coincide well with the two involved muscle groups. Further

evidence supporting the accuracy of this assignment is shown in FIG. 14. Plotted are time-series values of recovered absorption values at points in the image coinciding with the involved muscles. Noteworthy is the observation that the two signals are approximately 180° out of phase with each other, which is the expected response from the action of antagonistic muscle groups.

Imaging the Complexity of the Vascular Response.

One feature of the disclosed methods that may have considerable practical significance is the potential to obtain reliable estimates of the complexity of the vascular response. The significance of such measures is that they can reveal features of an integrated physiological state that are simply not measurable by using linear methods. Accordingly, in this study we have sought to confirm previous reports in the literature that indicate nonlinear chaotic behavior in vascular rhythms, as has been measured typically by surface laser Doppler methods. Specifically, we have computed certain measures that can reveal nonlinear behavior from time-series image data of the forearm of a subject performing a series of deep-breathing exercises. The influence of a respiratory stimulus was chosen as a simple noninvasive means to amplify the natural oscillatory activity of the vasculature. FIG. 15 shows an example of the magnitude of the signal response caused by deep breathing. The measures we have computed from the pixel data include ν , the correlation dimension, and, λ_1 , the largest Lyapunov exponent. Positive values for the latter indicate divergence of trajectories in state space, a finding consistent with a chaotic time series. A value of zero indicates no net divergence and is observed with stochastic data, while negative values indicate system damping.

The computed values displayed in a cross-sectional shown in FIG. 16, below were obtained using the method of Rosenstein *et al.* (Physica D, 65, 117-134 (1993)) for small data sets. The image series comprised 240 consecutive time points reconstructed from data collected at a rate of 2.8 Hz. The respiration rate was 0.08 Hz for the first 150
5 time points, and then doubled for the remainder of the measurement period. To assist in timing and repeatability, the volunteer was asked to adjust his respiratory rate to match the beat of a metronome.

Referring to FIG. 16, each pixel in the derived image series was band-pass filtered over a range that included the vasomotor and respiratory frequencies (0.05–0.35
10 Hz). Such filtering colors the noise and can lead to spurious low values for the correlation dimension. For this reason data obtained from a stochastic time series in a similar fashion was treated similarly. The control data set was obtained by computing an image time series from measurements performed on a solid white plastic rod composed of white Delrin®, whose scattering properties are similar to those of tissue. The rod diameter was
15 9 cm.

As shown in FIG. 16, the range of correlation dimensions computed from the physiological image time series is between 2 and 4 for most pixels. Significantly, these values are in close agreement with previous studies that measured the correlation dimensions of arterial and arteriolar vasomotion in *in vitro* rat or rabbit preparations. A
20 similar analysis on the rod data, shown in FIG. 17, reveals significantly higher values, a finding consistent with the character of the time series.

The image in FIG. 18 shows a spatial map of λ_1 computed for each pixel. Because a stochastic time series can also yield positive values for λ_1 , we have computed

surrogate data sets for each pixel and performed a test for statistical significance. Surrogate data were computed by randomizing the phase of the Fourier spectrum followed by recovery of the resultant time series from which the surrogate λ_1 were computed. FIG. 19 shows a map of the significance level, in each pixel, for the physiological data compared to surrogate data. Inspection reveals that most areas of the map have positive λ_1 values that fail the null hypothesis. Note that the optical measures performed are known to exhibit considerable sensitivity to the *microvessels*. These findings, indicating the occurrence of nonlinear chaotic activity of the vasculature revealed in a cross-sectional view, thus coincide well with previous reports whose measures were limited to the study of vascular dynamics in the near surface of tissue. Its worth emphasizing that the above findings are the first of their kind and, to the best of our knowledge, no other imaging modality is capable of providing equivalent measures of vascular dynamics.

Strategies of the types explored can be used to investigate a range of vascular anomalies. For example, the deep breathing exercise may aid in the diagnosis of cancer. Solid tumors have altered vascular beds. Enhanced breathing will cause redistribution of the flow patterns in tissue. Regions having altered vascular architecture should behave very different from those with a normal architecture. A similar approach can be used to identify tissue regions "at risk" due to the effects of diabetes, atherosclerosis (small vessel disease), or long term effects of smoking (large vessel disease).

Although the examples above focus on near infrared energy sources for imaging human tissue, the methodology of the present invention is applicable to essentially any wavelength for any energy source (e.g., electromagnetic, acoustic, etc.), responsive to

dynamic properties of a scattering medium and for any source condition (e.g., time independent, time harmonic, time resolved). Moreover, although the examples above investigate cross-sectional imaging, it will be appreciated that these methods are applicable to and include volumetric measures and imaging.

- 5 Further, although illustrative embodiments have been described herein in detail, those skilled in the art will appreciate that variations may be made without departing from the spirit and scope of this invention. Moreover, unless otherwise specifically stated, the terms and expressions used herein are terms of description and not terms of limitation, and are not intended to exclude any equivalents of the system and methods set
- 10 forth in the following claims.

What is claimed is:

1. A method for enhanced imaging of a target medium comprising:
directing energy into a target medium from at least one source during a period of time, the target medium having dynamic properties during the period of time;
measuring the density of the energy emerging from the target medium during the period of time using at least one detector, the density of the energy emerging from the target medium being a function of the dynamic properties of the target medium; and
generating a map of the dynamic properties of the target medium based on the measured density of energy emerging from the target medium.
2. The method of claim 1, further comprising:
generating a time series of images of the properties of a target medium based on the measured energy emerging from the target medium, wherein each image represents the cross-sectional properties of the target medium at a time interval during the period of time.
3. The method of claim 2, wherein generating the map of the dynamic properties of the target medium comprises processing the time series of images using time series analysis methods.
4. The method of claim 1, wherein generating the map of the dynamic properties of the target medium comprises processing the measured energy at each detector using time series analysis methods.

5. The method of claim 1, wherein the image of the dynamic properties is generated using time series analysis methods.

6. The method of claim 5, wherein the time series analysis methods comprise linear time series methods.

7. The method of claim 6, wherein the linear time series analysis methods are at least one of frequency analysis, time correlation analysis, time frequency analysis and principle component analysis.

8. The method of claim 5, wherein the time series analysis methods comprise nonlinear time series analysis methods.

9. The method of claim 1, wherein measuring the energy comprises a plurality of measurements collected at a sampling rate not less than twice the reciprocal of the highest frequency of a dynamic property to be imaged.

10. The method of claim 1, further comprising applying a provocation to the target medium.

11. The method of claim 1, wherein the target medium is human tissue having a vascular tree and the provocation has a dynamic effects on the vascular tree.

12. The method of claim 11, wherein the provocation is an autonomic stimulus.

13. The method of claim 11, wherein the provocation is a local stimulus.

14. The method of claim 1, wherein the energy is optical energy having a wavelength in the near infrared region of the electromagnetic spectrum.

15. The method of claim 14, wherein the optical energy directed toward the medium includes at least two wavelengths of near infrared energy.

16. The method of claim 14, wherein the target medium is human tissue having a vascular tree containing blood, the vascular tree comprising veins, arteries and micro vessels, the blood having time varying absorption and scattering properties to the near infrared energy as a function of blood oxygenation and blood volume.

17. The method of claim 16, wherein generating an image of the dynamic properties of the target medium comprises generating an image of at least one of the time varying absorption and scattering properties of the target medium.

18. The method of claim 16, wherein generating an image of the dynamic properties of the target medium further comprises using time series analysis to enhance the contrast of at least one of veins, arteries and micro vessels.

19. The method of claim 1, wherein the energy is optical energy in the visible spectrum.

20. The method of claim 1, wherein generating an image of the dynamic properties of the target medium further comprises processing the measured energy using a modified perturbation formulation of the radiation transport equation, wherein the modified perturbation formulation uses relative energy measurements.

21. The method of claim 20, wherein the relative energy measurements are the relative differences between a measure at an instant in time and a time average mean of measures during the period of time.

22. A system for enhanced imaging of a target medium comprising:

a source for directing energy into a target medium from at least one source during a period of time, the target medium having dynamic properties during the period of time;

a detector for measuring the density of the energy emerging from the target medium during the period of time using at least one detector, the density of the energy emerging from the target medium being a function of the dynamic properties of the target medium;

a data acquisition means for receiving the measured energy densities during the period of time; and

a computer connected to the data acquisition means, the computer having code for generating a map of the dynamic properties of the target medium based on the measured density of energy emerging from the target medium.

23. The system of claim 22, wherein the computer further includes code for generating a time series of images of the properties of a target medium based on the measured energy emerging from the target medium, wherein each image represents the cross-sectional properties of the target medium at a time interval during the period of time.

24. The system of claim 23, wherein the computer further includes code for processing the time series of images using time series analysis methods to generate the map of the dynamic properties.

25. The system of claim 24, wherein the computer further includes code for processing the measured energy at each detector using time series analysis methods to generate the map of the dynamic properties of the medium.

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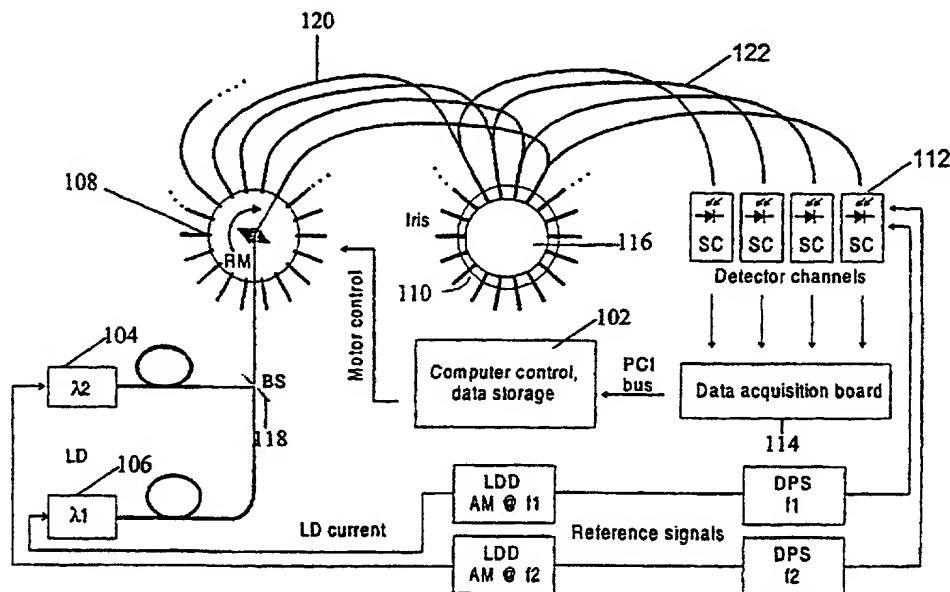
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(54) Title: METHOD AND SYSTEM FOR IMAGING THE DYNAMICS OF SCATTERING MEDIUM



(57) Abstract: A method and system for imaging the dynamics of a scattering medium (116) is provided. The method and system generates contrast and resolution enhanced images of dynamic properties of a medium having a temporal signature by using time series analysis methods on a time series of collected data or time series of images to extract and isolate dynamic properties of the medium (116).

WO 01/20305 A1

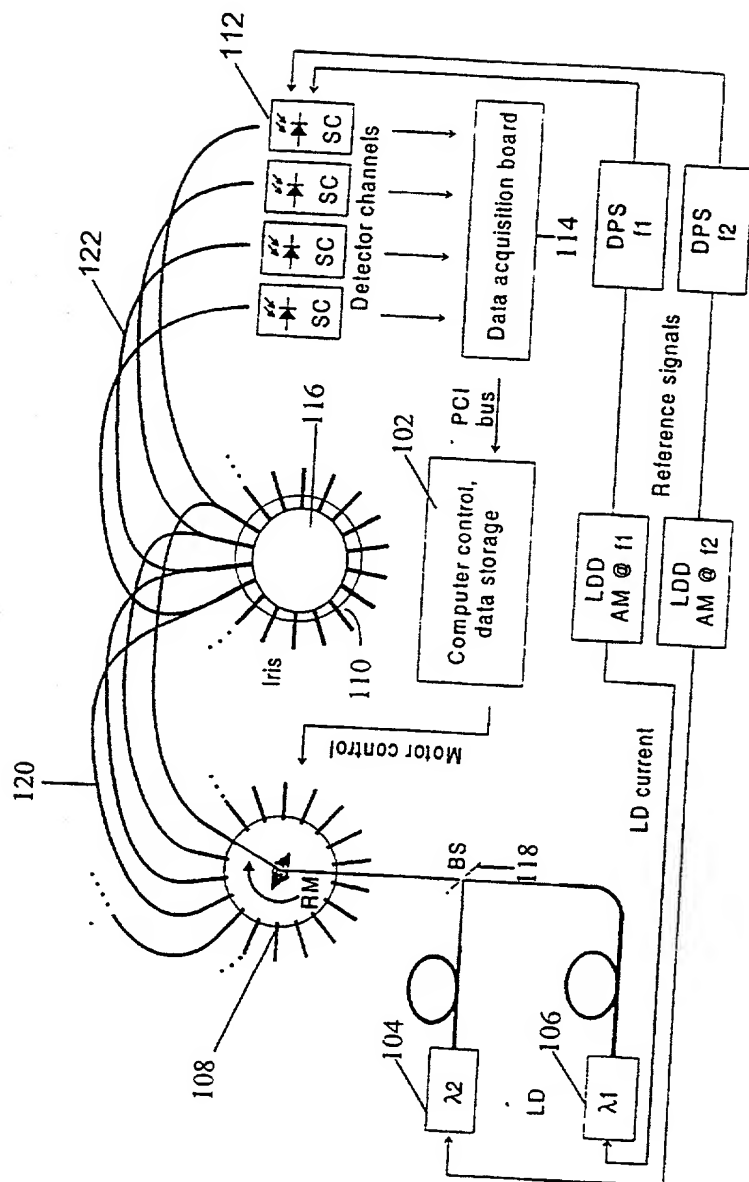


FIG. 1

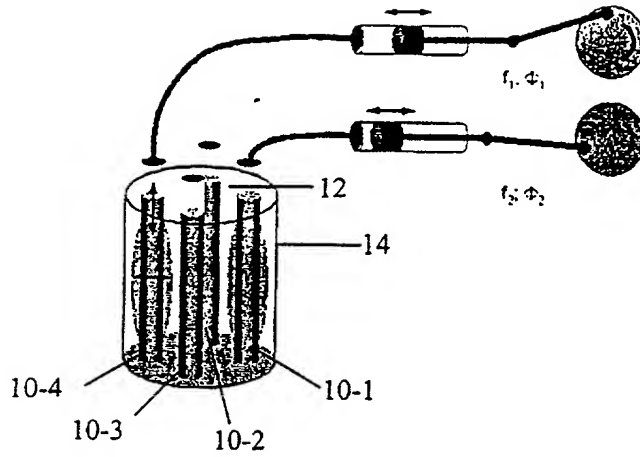


Fig. 2A

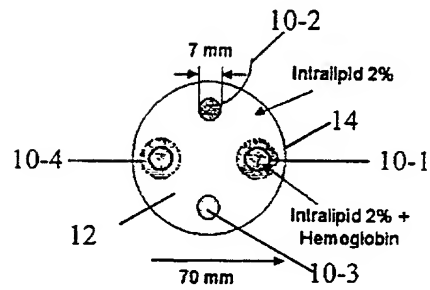


Fig. 2B

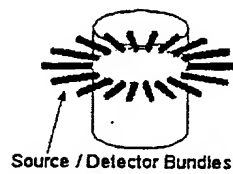


Fig. 2C

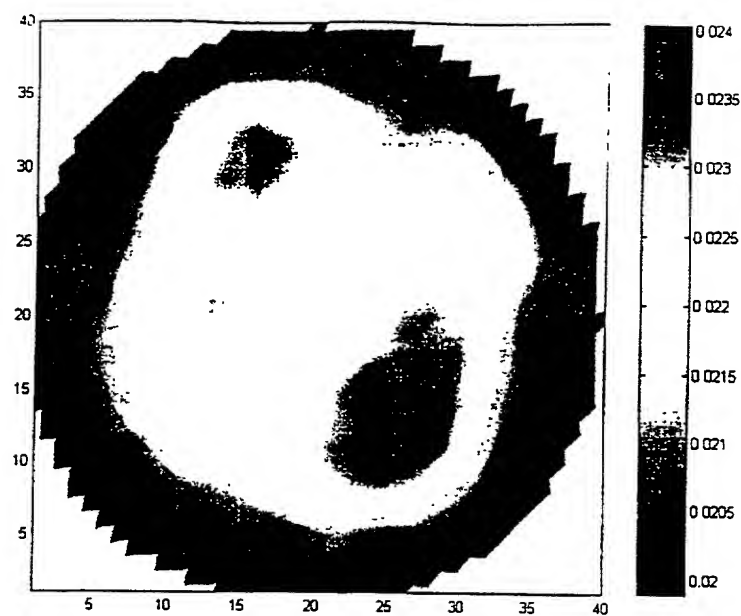


Fig. 3

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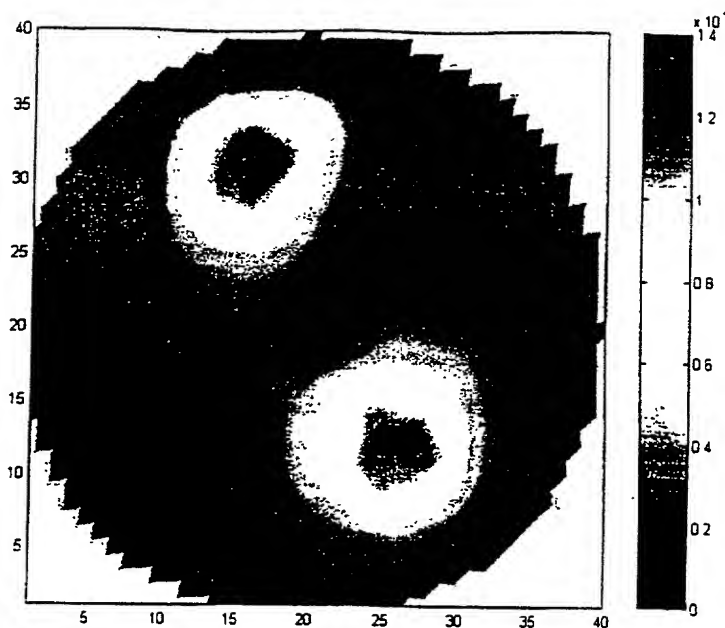


Fig. 4

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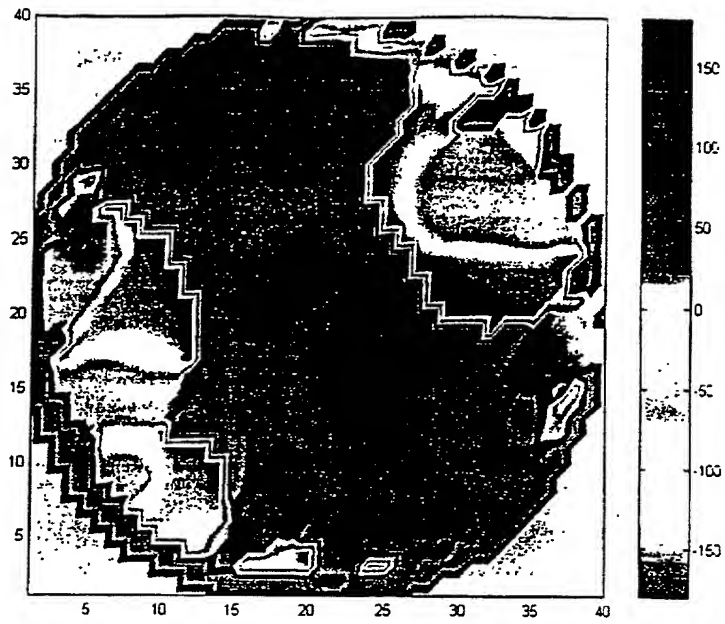


Fig. 5

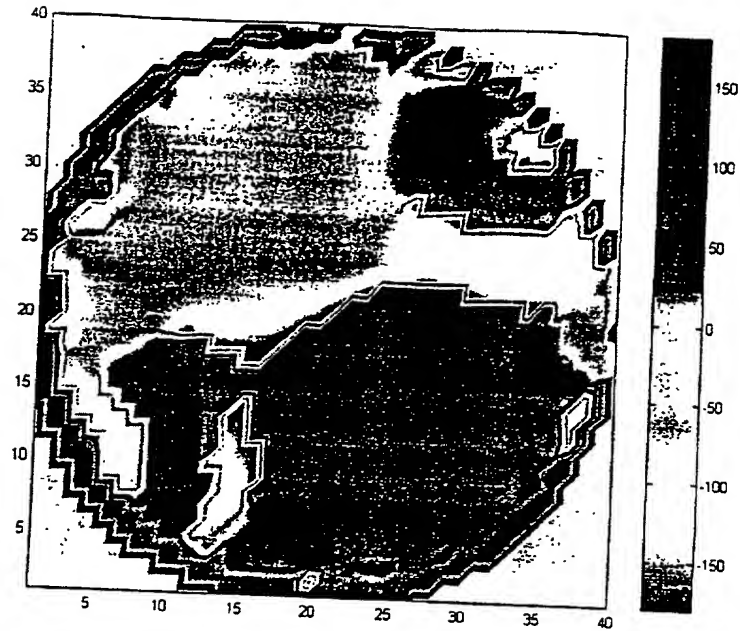


Fig. 6

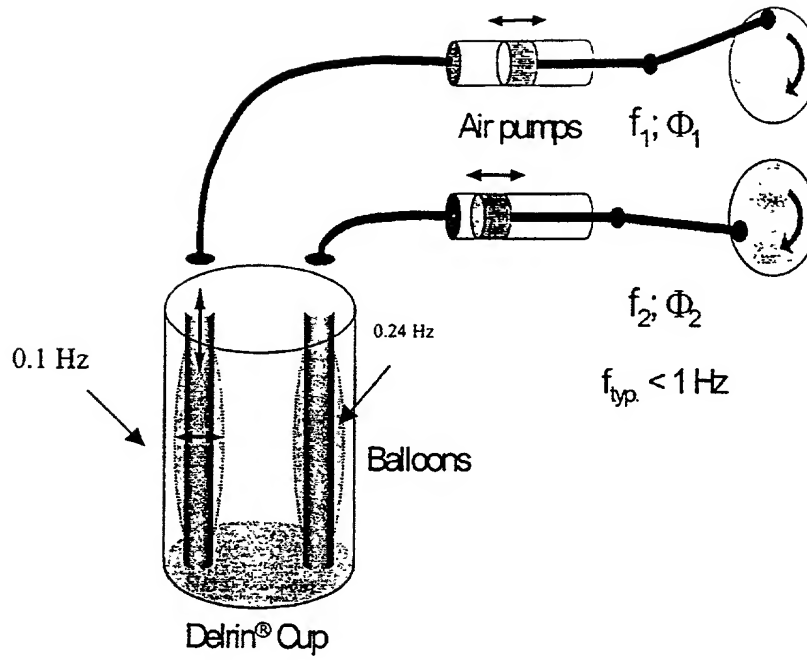


Fig. 7A

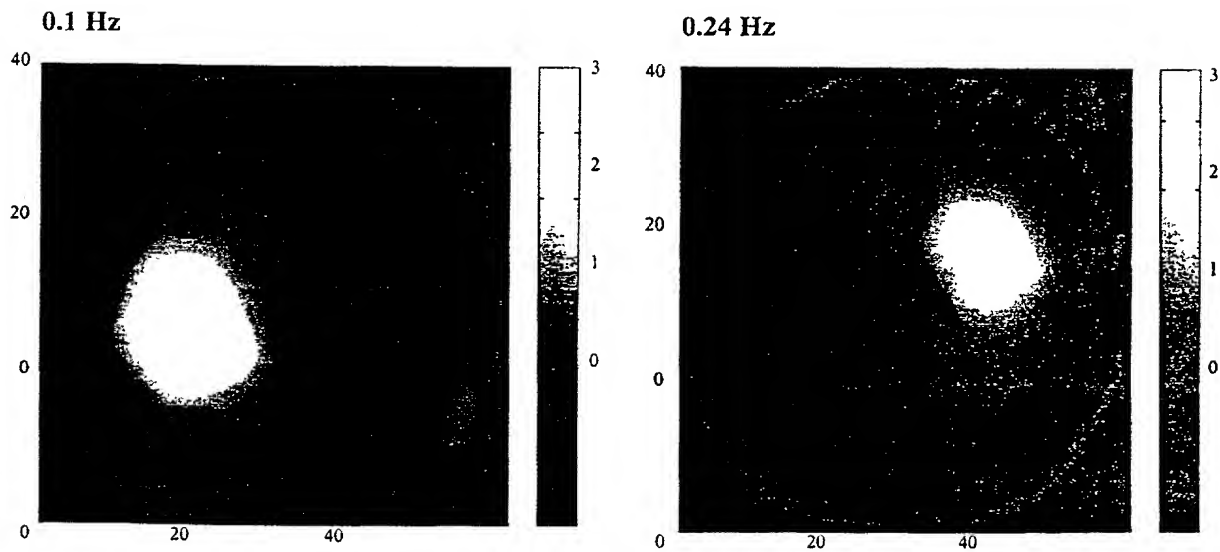


Fig. 7B

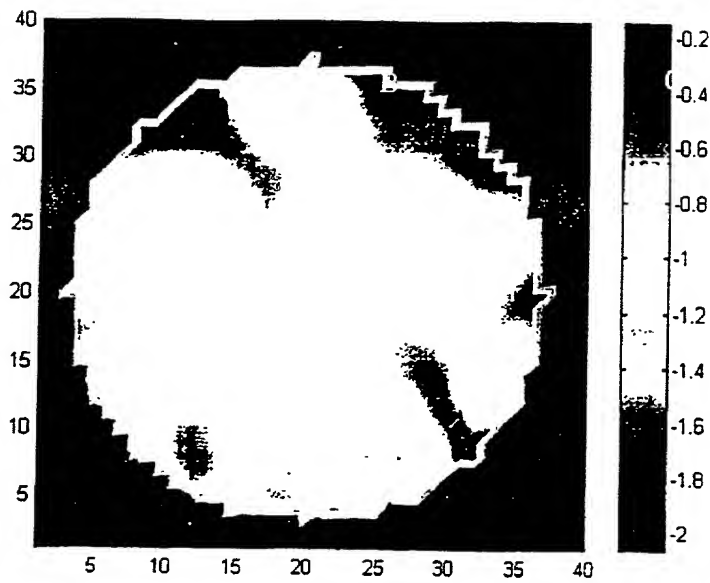


Fig. 8



Fig. 9



Fig. 10

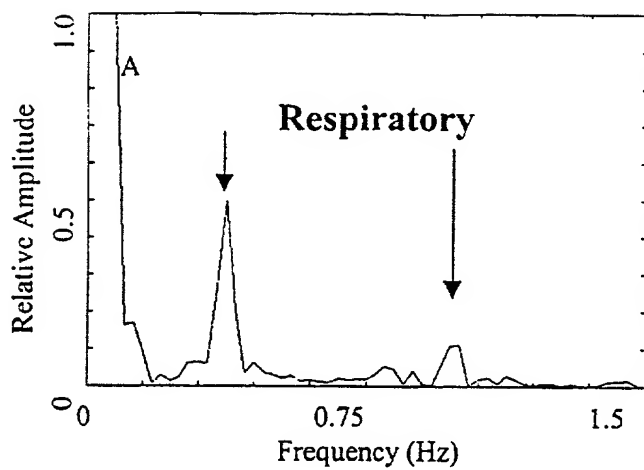


Fig. 11A

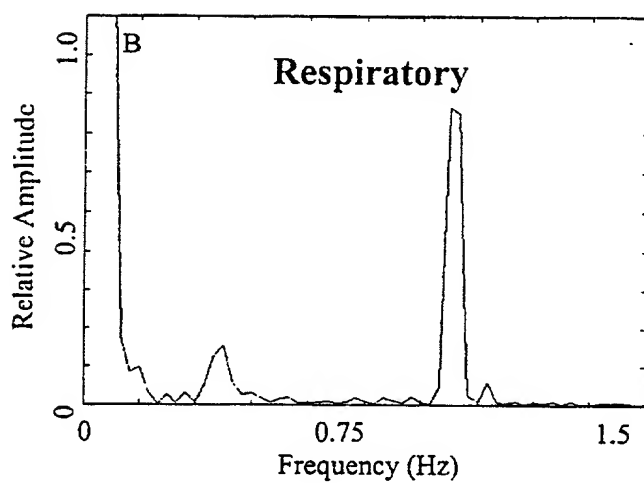


Fig. 11B

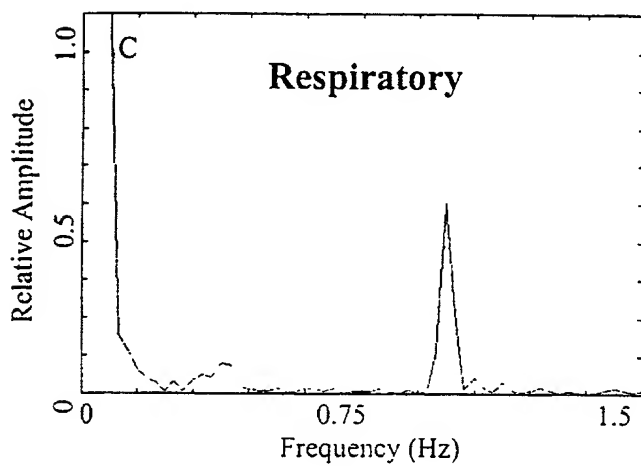


Fig. 11C

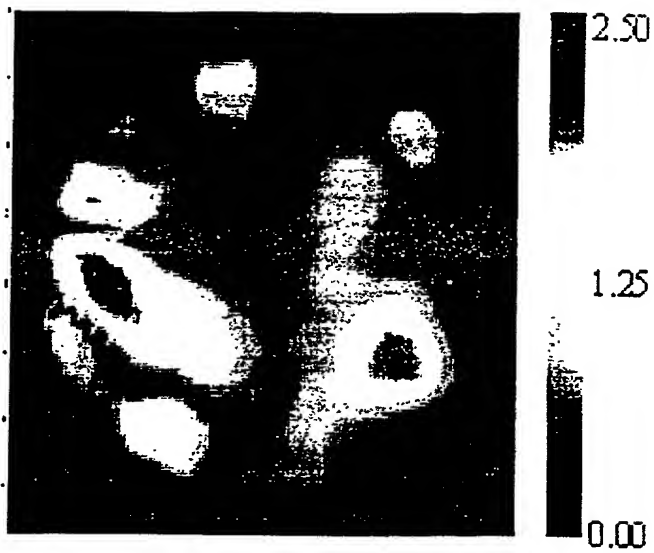


Fig.12

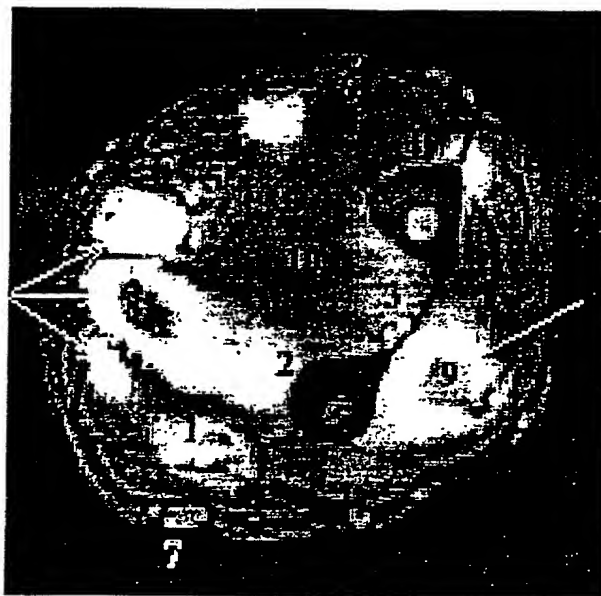


Fig. 13

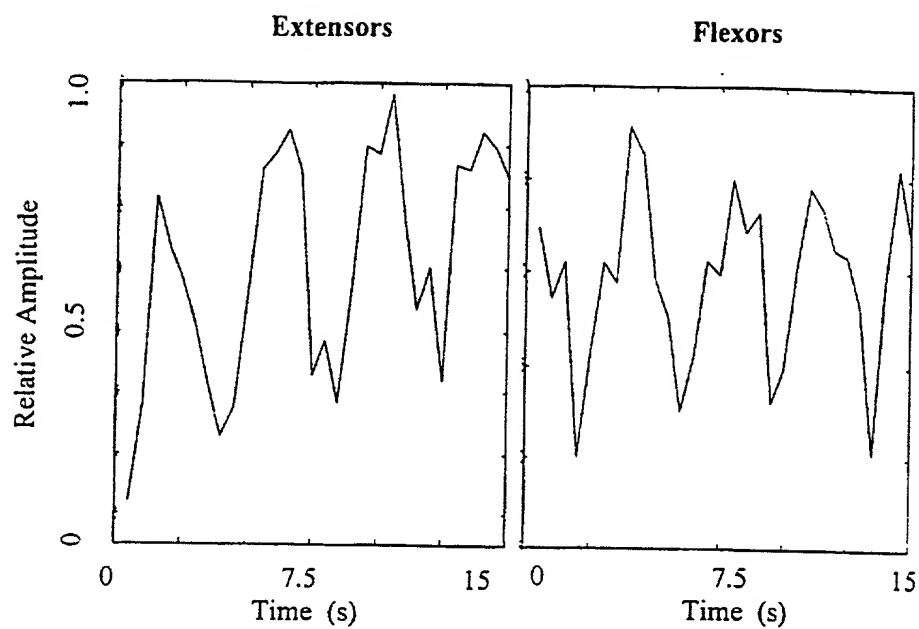


Fig. 14

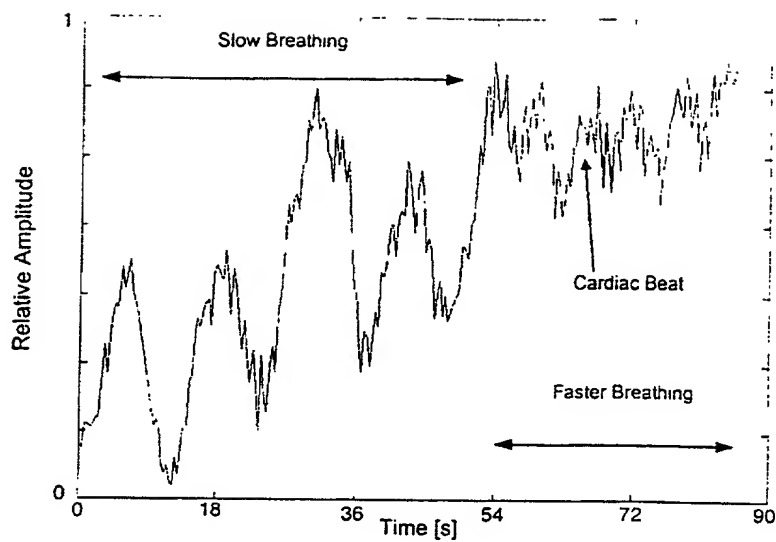


Fig. 15

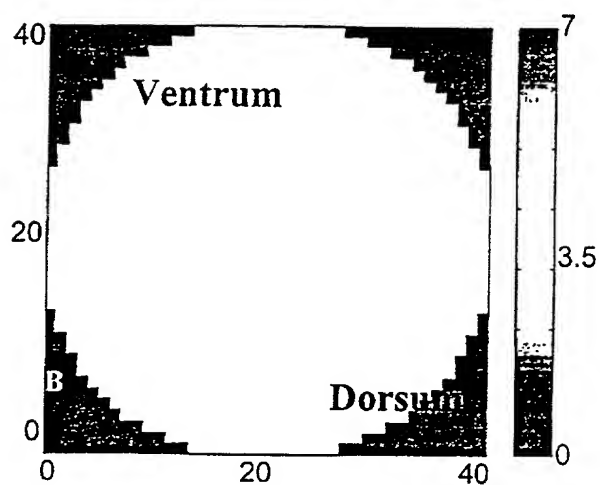


Fig. 16

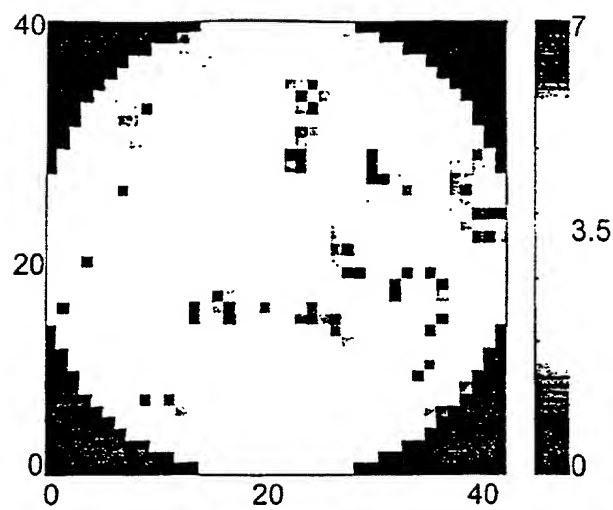


Fig. 17

90/000000

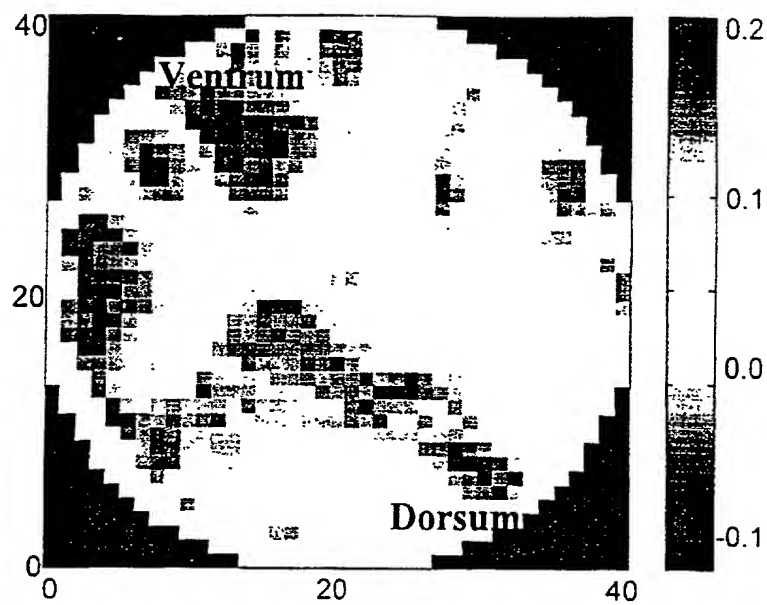


Fig. 18

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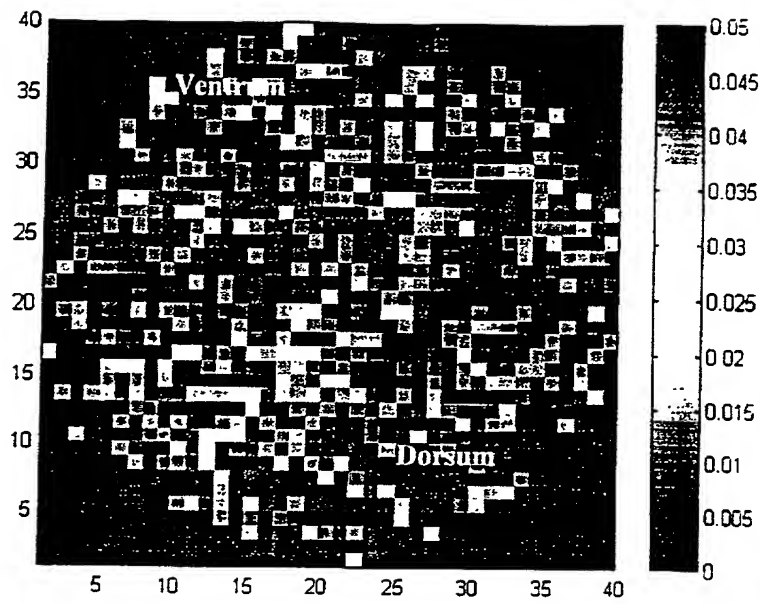


Fig. 19

COMBINED DECLARATION AND POWER OF ATTORNEY FOR
ORIGINAL, DESIGN, NATIONAL STAGE OF PCT, SUPPLEMENTAL
DIVISIONAL, CONTINUATION OR CONTINUATION-IN-PART APPLICATION

As a below name inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

METHOD AND SYSTEM FOR IMAGING THE DYNAMICS OF A SCATTERING MEDIUM
the specification of which

- a. ☐ is attached hereto
- b. ☐ was filed on _____ as application Serial No. _____ and was amended on _____ (if applicable).

PCT FILED APPLICATION ENTERING NATIONAL STAGE

- c. ☒ was described and claimed in International Application No. PCT/US00/25136 filed on 14 September 2000 and as amended on _____ (if any).

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the patentability as defined in Title 37, Code of Federal Regulations, § 1.56.

I hereby specify the following as the correspondence address to which all communications about this application are to be directed:

SEND CORRESPONDENCE TO: MORGAN & FINNEGAN, L.L.P
345 Park Avenue
New York, N.Y. 10154

DIRECT TELEPHONE CALLS TO: Tod M. Melgar
(212) 758-4800

☐ I hereby claim foreign priority benefits under Title 35, United States Code § 119(a)-(d) or under § 365(b) of any foreign application(s) for patent or inventor's certificate or under § 365(a) of any PCT international application(s) designating at least one country other than the U.S. listed below and also have identified below such foreign application(s) for patent or inventor's certificate or such PCT international application(s) filed by me on the same subject matter having a filing date within twelve (12) months before that of the application on which priority is claimed:

☐ The attached 35 U.S.C. § 119 claim for priority for the application(s) listed below forms a part of this declaration.

<u>Country/PCT</u>	<u>Application Number</u>	<u>Date of filing (day, month, yr)</u>	<u>Date of Issue (day, month, yr)</u>	<u>Priority Claimed</u>
				<input type="checkbox"/> YES <input type="checkbox"/> NO
				<input type="checkbox"/> YES <input type="checkbox"/> NO
				<input type="checkbox"/> YES <input type="checkbox"/> NO

[X] I hereby claim the benefit under 35 U.S.C. § 119(e) of any U.S. provisional application(s) listed below.

<u>Provisional Application No.</u>	<u>Date of Filing (day, month, yr)</u>
60/153,926	14 September 1999
60/154,099	15 September 1999

ADDITIONAL STATEMENTS FOR DIVISIONAL, CONTINUATION OR CONTINUATION-IN-PART OR PCT INTERNATIONAL APPLICATION(S) (DESIGNATING THE U.S.)

I hereby claim the benefit under Title 35, United States Code § 120 of any United States application(s) or under § 365(c) of any PCT international application(s) designating the U.S. listed below.

<u>US/PCT Application Serial No.</u>	<u>Filing Date</u>	<u>Status (patented, pending, abandoned)/ U.S. application no. assigned (For PCT)</u>

☐ In this continuation-in-part application, insofar as the subject matter of any of the claims of this application is not disclosed in the above listed prior United States or PCT international application(s) in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, § 1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or Imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

35 I hereby appoint the following attorneys and/or agents with full power of substitution and revocation, to prosecute this application, to receive the patent, and to transact all business in the Patent and Trademark Office connected therewith: John A. Diaz (Reg. No. 19,550), John C. Vassil (Reg. No. 19,098), Alfred P. Ewert (Reg. No. 19,887), David H. Pfeffer (Reg. No. 19,825), Harry C. Marcus (Reg. No. 22,390), Robert E. Paulson (Reg. No. 21,046), Stephen R. Smith (Reg. No. 22,615), Kurt E. Richter (Reg. No. 24,052), J. Robert Dailey (Reg. No. 27,434), Eugene Moroz (Reg. No. 25,237), John F. Sweeney (Reg. No. 27,471), Arnold I. Rady (Reg. No. 26,601), Christopher A. Hughes (Reg. No. 26,914), William S. Feiler (Reg. No. 26,728), Joseph A. Calvaruso (Reg. No. 28,287), James W. Gould (Reg. No. 28,859), Richard C. Komson (Reg. No. 27,913), Israel Blum (Reg. No. 26,710), Bartholomew Verdirame (Reg. No. 28,483), Maria C.H. Lin (Reg. No. 29,323), Joseph A. DeGirolamo (Reg. No. 28,595), Michael P. Dougherty (Reg. No. 32,730), Seth J. Atlas (Reg. No. 32,454), Andrew M. Riddles (Reg. No. 31,657), Bruce D. DeRenzi (Reg. No. 33,676), Michael M. Murray (Reg. No. 32,537), Mark J. Abate (Reg. No. 32,527), John T. Gallagher (Reg. No. 35,516), Steven F. Meyer (Reg. No. 35,613), Kenneth H. Sonnenfeld (Reg. No. 33,285), Tony V. Pezzano (Reg. No. 38,271), Andrea L. Wayda (Reg. No. 43,979) and Walter G. Hanchuk Reg. No. (35,179) of Morgan & Finnegan, L.L.P. whose address is: 345 Park Avenue, New York, New York, 10154; and Michael S. Marcus (Reg. No. 31,727) and John E. Hoel (Reg. No. 26,279) of Morgan & Finnegan, L.L.P., whose address is 1775 Eye Street, Suite 400, Washington, D.C. 20006.

[] I hereby authorize the U.S. attorneys and/or agents named hereinabove to accept and follow instructions from _____ as to any action to be taken in the U.S. Patent and Trademark Office regarding this application without direct communication between the U.S. attorneys and/or agents and me. In the event of a change in the person(s) from whom instructions may be taken I will so notify the U.S. attorneys and/or agents hereinabove.

Full name of sole or first inventor Randall L. BARBOUR

Inventor's signature* Randall L. Barbour

11/14/2000
date

Residence 15 Cherry Lane, Glen Head, New York 11546 NY

Citizenship United States

Post Office Address same as above

Full name of second joint inventor, if any _____

Inventor's signature* _____

date

Residence _____

Citizenship _____

Post Office Address _____

Full name of third joint inventor, if any _____

Inventor's signature* _____ date _____

Residence _____

Citizenship _____

Post Office Address _____

[] ATTACHED IS/ARE ADDED PAGE(S) TO COMBINED DECLARATION AND POWER OF ATTORNEY FORM FOR SIGNATURE BY FOURTH AND SUBSEQUENT INVENTORS

* Before signing this declaration, each person signing must:

1. Review the declaration and verify the correctness of all information therein; and
2. Review the specification and the claims, including any amendments made to the claims.

After the declaration is signed, the specification and claims are not to be altered.

To the inventor(s):

The following are cited in or pertinent to the declaration attached to the accompanying application:

Title 37, Code of Federal Regulation, § 1.56

Duty to disclose information material to patentability.

(a) A patent by its very nature is affect with a public interest. The public interest is best served, and the most effective patent examination occurs when, at the time an application is being examined, the Office is aware of and evaluates the teachings of all information material to patentability. Each individual associated with the filing and prosecution of a patent application has a duty of candor and good faith in dealing with the Office, which includes a duty to disclose to the Office all information known to that individual to be material to patentability as defined in this section. The duty to disclose information exists with respect to each pending claim until the claim is canceled or withdrawn from consideration, or the application becomes abandoned. Information material to the patentability of a claim that is canceled or withdrawn from consideration need not be submitted if the information is not material to the patentability of any claim remaining under consideration in the application. There is no duty to submit information which is not material to the patentability of any existing claim. The duty to disclose all information known to be material to patentability is deemed to be satisfied if all information known to be material to patentability of any claim issued in patent was cited by the Office or submitted to the Office in the manner prescribed by §§1.97(b)-(d) and 1.98. However, no patent will be granted on an application in connection with which fraud on the Office was practiced or attempted or the duty of disclosure was violated through bad faith or intentional misconduct. The Office encourages applicants to carefully examine:

- (1) prior art cited in search reports of a foreign patent office in a counterpart application, and

- (2) the closest information over which individuals associated with the filing or prosecution of a patent application believe any pending claim patentably defines, to make sure that any material information contained therein is disclosed to the Office.

Title 35, U.S. Code § 101

Inventions patentable

Whoever invents or discovers any new and useful process, machine, manufacture, or composition of matter, or any new and useful improvement thereof, may obtain a patent therefor, subject to the conditions and requirements of this title.

Title 35 U.S. Code § 102

Conditions for patentability; novelty and loss of right to patent

A person shall be entitled to a patent unless –

- (a) the invention was known or used by others in this country, or patented or described in a printed publication in this or a foreign country, before the invention thereof by the applicant for patent,
- (b) the invention was patented or described in a printed publication in this or foreign country or in public use or on sale in this country, more than one year prior to the date of application for patent in the United States, or
- (c) he has abandoned the invention, or
- (d) the invention was first patented or caused to be patented, or was the subject of an inventor's certificate, by the applicant or his legal representatives or assigns in a foreign country prior to the date of the application for patent in this country on an application for patent or inventor's certificate filed more than twelve months before the filing of the application in the United States, or
- (e) the invention was described in a patent granted on an application for patent by another filed in the United States before the invention thereof by the applicant for patent, or on an international application by another who has fulfilled the requirements of paragraphs (1), (2), and (4) of section 371(c) of this title before the invention thereof by the applicant for patent, or
- (f) he did not himself invent the subject matter sought to be patented, or
- (g) before the applicant's invention thereof the invention was made in this country by another had not abandoned, suppressed, or concealed it. In determining priority of invention there shall be considered not only the respective dates of conception and reduction to practice of the invention, but also the reasonable diligence of one who was first to conceive and last to reduce to practice, from a time prior to conception by the other ...

Title 35, U.S. Code § 103

Conditions for patentability; non-obvious subject matter

A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having

ordinary skill in the art to which said matter pertains. Patentability shall not be negated by the manner in which the invention was made.

Subject matter developed by another person, which qualifies as prior art only under subsection (f) or (g) of section 102 of this title, shall not preclude patentability under this section where the subject matter and the claimed invention were, at the time the invention was made, owned by the same person or subject to an obligation of assignment to the same person.

Title 35, U.S. Code § 112 (in part)

Specification

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise and exact terms also enable any person skilled in the art to which it pertains, or with which it is mostly nearly connected, to make and use the same, and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Title 35, U.S. Code § 119

Benefit of earlier filing date in foreign country; right of priority

An application for patent for an invention filed in this country by any person who has, or whose legal representatives or assigns have, previously regularly filed an application for a patent for the same invention in a foreign country which affords similar privileges in the case of applications filed in the United States or to citizens of the United States, shall have the same effect as the same application would have if filed in this country on the date on which the application for patent for the same invention was first filed in such foreign country, if the application in this country is filed within twelve months from the earliest date on which such foreign application was filed; but no patent shall be granted on any application for patent for an invention which had been patented or described in a printed publication in any country more than one year before the date of the actual filing of the application in this country, or which had been in public use or on sale in this country more than one year prior to such filing.

Title 35, U.S. Code § 120

Benefit or earlier filing date in the United States

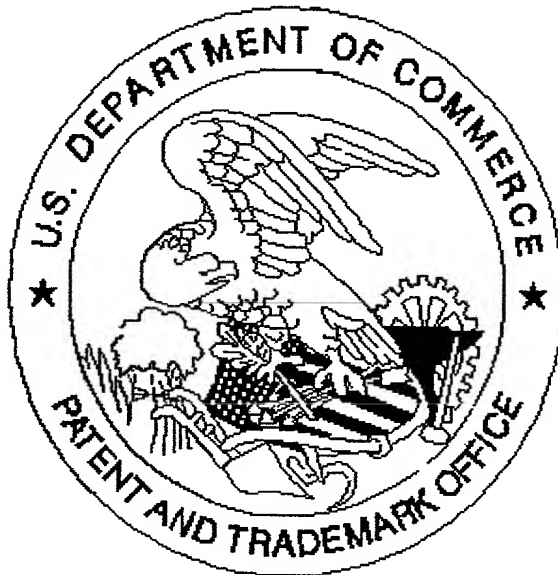
An application for patent for an invention disclosed in the manner provided by the first paragraph of section 112 of this title in an application previously filed in the United States, or as provided by section 363 of this title, which is filed by an inventor or inventors named in the previously filed application shall have the same effect, as to such invention, as though filed on the date of the prior application, if filed before the patenting or abandonment of or termination of proceedings on the first application or an application similarly entitled to the benefit of the filing date of the first application and if it contains or is amended to contain a specific reference to the earlier filed application.

Please read carefully before signing the Declaration attached to the accompanying Application.

If you have any questions, please contact Morgan & Finnegan, L.L.P.

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Rev. 10/00

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Application deficiencies found during scanning:

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